

## **PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix**

Catalog Numbers A25741, A25742, A25743, A25776, A25777, A25778, A25779, A25780, A25918

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## **Product Information**

#### About the Reagent

The PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix, is formulated to provide superior specificity and sensitivity. It is supplied in a convenient 2X concentration premix to perform real-time PCR using SYBR<sup>™</sup> Green dye. The master mix contains:

- SYBR<sup>™</sup> Green Dye
- Dual-Lock<sup>™</sup> DNA Polymerase, with a proprietary combination of two proprietary hot start modifications for exceptional specificity
- Heat-labile Uracil-DNA Glycosylase (UDG)
- ROX<sup>™</sup> dye Passive Reference
- dNTP blend containing dUTP/dTTP
- Optimized buffer components

The user only needs to provide primers, template, and water.

#### **Hot Start**

PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix contains the proprietary Dual-Lock<sup>™</sup> *Taq* DNA Polymerase that utilizes a combination of two hot start mechanisms to control its activity.

The dual hot start approach provides tight control over *Taq* activation, preventing undesirable early activity of the polymerase at low temperatures that can lead to non-specific amplification.

It allows users flexibility in reaction set up, including the pre-mixing of PCR reagents and storage at room temperature for up to 72 hours prior to cycling.

The polymerase is reactivated after only a 2 minute incubation at 95°C.

#### UDG

PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix contains heat-labile uracil-DNA glycosylase (UDG). UDG is also known as uracil-N-glycosylase (UNG).

Treatment with heat-labile UDG can prevent the reamplification of carryover PCR products by removing any uracil incorporated into single- or double-stranded amplicons (Longo et al., 1990). Heat-labile UDG prevents reamplification of carryover PCR products in an assay if all previous PCR for that assay was performed using a dUTP-containing master mix. See "Prevent Contamination and Nonspecific Amplification" on page 11 for more information about UDG.

PCR products are stable for up to 72 hours post-amplification using master mixes containing heat-labile UDG. Unlike standard UDG, heat-labile UDG is completely inactivated prior to amplification.

#### dUTP/dTTP

A blend of dUTP/dTTP is included to enable UDG activity and maintain optimal PCR results.

#### SYBR™ Green

The SYBR™ Green dye detects PCR products by binding to double stranded DNA formed during PCR (see Chemistry Overview section).

#### ROX Passive Reference

PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix contains  $ROX^{™}$  dye Passive Reference. The  $ROX^{™}$  dye Passive Reference provides an internal reference to which the reporter-dye signal can be normalized during data analysis. Normalization is necessary to correct for fluorescence fluctuations due to changes in concentration or volume.

#### Real-Time Instruments

Power $Up^{\scriptscriptstyle{TM}}$  SYBR $^{\scriptscriptstyle{TM}}$  Green Master Mix can be used to run experiments on the following Applied Biosystems Real-Time PCR Systems:

- StepOne<sup>™</sup> and StepOnePlus<sup>™</sup> Real-Time PCR Systems
- 7500 and 7500 Fast Real-Time PCR Systems
- 7900HT and 7900HT Fast Real-Time PCR Systems
- ViiA<sup>™</sup> 7 Real-Time PCR System
- QuantStudio<sup>™</sup> family of Real-Time PCR Systems

## About This Protocol

#### This protocol provides:

- Background information about gene quantification assays
- A list of equipment and materials for using the PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix
- Procedures for using the PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix

For details about specific procedures described in this protocol, see "Support Documents" on page 33.

### **Chemistry Overview**

How the SYBR™ Green Dye Chemistry Works The SYBR™ Green dye is used to detect PCR products by binding to double-stranded DNA formed during PCR. The process works as follows:

- 1. When PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix is added to a sample, SYBR<sup>™</sup> Green dye immediately binds to all double-stranded DNA.
- 2. During the PCR, Dual-Lock™ DNA Polymerase amplifies the target sequence, which creates the PCR product, or "amplicon."
- 3. The SYBR™ Green dye then binds to each new copy of double-stranded DNA.
- 4. As the PCR progresses, more amplicon is created. Because the SYBR™ Green dye binds to all double-stranded DNA, the result is an increase in fluorescence intensity proportional to the amount of double-stranded PCR product produced.

The following figure illustrates this process.

Step 1



The SYBR™ Green dye within the PowerUp™ SYBR™ Green Master Mix immediately binds with all double-stranded DNA

Step 2



During PCR, Dual-Lock™ DNA Polymerase amplifies each target.

Step 3



The SYBR™ Green dye then binds to each new copy of doublestranded DNA.

Figure 1 Representation of how the SYBR™ Green dye acts on double-stranded DNA during one extension phase of PCR

Using the Master Mix in Two-Step RT-PCR When performing a two-step RT-PCR reaction, total or mRNA must first be transcribed into cDNA:

- 1. In the reverse transcription (RT) step, cDNA is reverse transcribed from total RNA samples using random primers from the High-Capacity cDNA Reverse Transcription Kit or SuperScript $^{\text{TM}}$  VILO $^{\text{TM}}$  cDNA Synthesis Kit (see page 10).
- 2. In the PCR step, PCR products are synthesized from cDNA samples using the PowerUp $^{\text{\tiny TM}}$  SYBR $^{\text{\tiny TM}}$  Green Master Mix.

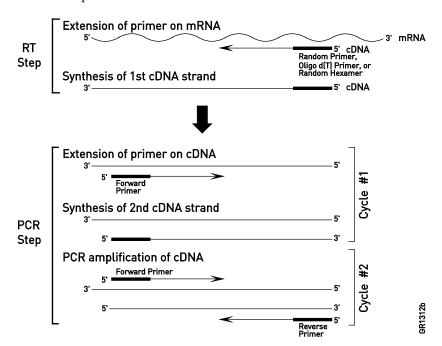


Figure 2 Two-step RT-PCR

## **Contents and Storage**

 $\textbf{Contents} \hspace{1cm} \textbf{The PowerUp}^{\text{\tiny{IM}}} \, \textbf{SYBR}^{\text{\tiny{IM}}} \, \textbf{Green Master Mix is supplied in a 2X concentration}.$ 

Catalog no.	Contents
A25741	PowerUp™ SYBR™ Green Master Mix, 1 mL
A25779	PowerUp™ SYBR™ Green Master Mix, 2 × 1 mL
A25780	PowerUp™ SYBR™ Green Master Mix, 5 × 1 mL
A25918	PowerUp™ SYBR™ Green Master Mix, 10 × 1 mL
A25742	PowerUp™ SYBR™ Green Master Mix, 5 mL
A25776	PowerUp™ SYBR™ Green Master Mix, 2 × 5 mL
A25777	PowerUp™ SYBR™ Green Master Mix, 5 × 5 mL
A25778	PowerUp™ SYBR™ Green Master Mix, 10 × 5 mL
A25743	PowerUp™ SYBR™ Green Master Mix, 50 mL

Storage Store the PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix at  $2^{\circ}$ C to  $8^{\circ}$ C.

## **Required Materials**

#### **Plates**

Choose the plate appropriate for your real-time instrument.

Instrument	Plates <sup>‡</sup>	Catalog number
	MicroAmp™ Fast Optical 96-Well Reaction Plate with Barcode (0.1-mL)	
	20 plates	4346906
	• 200 plates	4366932
ViiA™ 7 and	MicroAmp™ Fast Optical 96-Well Reaction Plate with Barcode	
QuantStudio™	• 20 plates	4306737
systems	• 500 plates	4326659
	MicroAmp™ Optical 384-Well Reaction Plate with Barcode	
	• 50 plates	4309849
	• 500 plates	4326270
	• 1000 plates	4343814
Step0ne <sup>™</sup> system	MicroAmp™ Fast Optical 48-Well Reaction Plate, 20 plates	4375816
StepOnePlus <sup>™</sup> and	MicroAmp™ Fast Optical 96-Well Reaction Plate with Barcode	
7500 Fast systems	(0.1-mL)	
	• 20 plates	4346906
	• 200 plates	4366932
7900HT and	MicroAmp™ Optical 96-Well Reaction Plate with Barcode	
7900HT Fast§	• 20 plates	4306737
systems	• 500 plates	4326659
	MicroAmp™ Optical 384-Well Reaction Plate with Barcode	
	• 50 plates	4309849
	• 500 plates	4326270
	• 1000 platesv	4343814
7500 system	MicroAmp™ Optical 96-Well Reaction Plate with Barcode	
-	• 20 plates	4306737
	• 500 plates	4326659

 $<sup>\</sup>S$  Requires a MicroAmp<sup>TM</sup> Snap-On Optical Film Compression Pad (Cat. no. 4333292) when using the standard MicroAmp<sup>TM</sup> Optical 96-Well Reaction Plate.

#### **Optical Seals**

Seal all plates, except StepOne $^{^{\text{\tiny TM}}}$  system plates, with MicroAmp $^{^{\text{\tiny TM}}}$  Optical Adhesive Film. Seal StepOne $^{^{\text{\tiny TM}}}$  system plates with MicroAmp $^{^{\text{\tiny TM}}}$  48-Well Optical Adhesive Film.

Item	Catalog number
MicroAmp™ 48-Well Optical Adhesive Film:	
25 covers	4375928
• 100 covers	4375323
MicroAmp™ Optical Adhesive Film:	
25 covers	4360954
• 100 covers	4311971

#### Other Kits

Item	Catalog number
High Capacity cDNA Reverse Transcription Kit:	
• 200 reactions	4368814
<ul> <li>200 reactions with RNase Inhibitor</li> </ul>	4374966
• 1000 reactions	4368813
• 1000 reactions with RNase Inhibitor	4374967
High-Capacity RNA-to-cDNA™ Kit	
• 50 reactions	4387406
SuperScript™ VILO™ cDNA Synthesis Kit:	
• 50 reactions	4453650
• 250 reactions	4453651

#### Other Consumables

**IMPORTANT**: Do not use plastics made of polyethylene terephthalate copolyester, glycol modified (PTEG) for storage of PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix or reaction mixes. SYBR<sup>™</sup> dye has been shown to stick to this type of plastic material.

Plastics recommended for storage include polypropylene, high density polyethylene (HDPE), and polystyrene.

Item	Source
Centrifuge with adapter for 96-well plates	Major laboratory supplier (MLS)
or	
Centrifuge with adapter for 384-well plates	
Disposable gloves	MLS
Microcentrifuge	MLS
Pipette tips, with filter plugs	MLS
Pipettors, positive-displacement or air-displacement	MLS
Polypropylene tubes	MLS
Tris-EDTA (TE) Buffer, pH 8.0	MLS
Vortexer	MLS

### Prevent Contamination and Nonspecific Amplification

#### **Overview**

PCR assays require special laboratory practices to avoid false positive amplifications. The high throughput and repetition of these assays can lead to amplification of a single DNA molecule.

#### Using UDG to Minimize Reamplification Carryover Products

PowerUp™ SYBR™ Green Master Mix contains heat-labile uracil-DNA glycosylase (UDG). UDG is also known as uracil-N-glycosylase (UNG). Treatment with heat-labile UDG is useful in preventing the reamplification of carryover PCR products.

The heat-labile UDG used in the PowerUp<sup>TM</sup> SYBR<sup>TM</sup> Green Master Mix is a 26-kDa recombinant enzyme derived from the thermolabile UDG gene isolated from marine bacteria, and expressed in *E. coli*.

UDG acts on single- and double-stranded dU-containing DNA. It acts by hydrolyzing uracil-glycosidic bonds at dU-containing DNA sites. The enzyme causes the release of uracil, thereby creating an alkali-sensitive apyrimidic site in the DNA. The enzyme has no activity on RNA or dT-containing DNA (Longo et al., 1990).

#### Using NTC Controls

No Template Control (NTC) reactions can be used to identify PCR contamination. NTC reactions contain all reaction components (PowerUp $^{\text{\tiny TM}}$  SYBR $^{\text{\tiny TM}}$  Green Master Mix, primers, water) except sample, and therefore should not return a  $C_T$  value.

#### Design Primers to Avoid Primer-Dimers

Use primers that contain dA nucleotides near the 3´ ends so that any primer-dimer generated is efficiently degraded by UDG at least as well as any dU-containing PCR products. The farther a dA nucleotide is from the 3´ end, the more likely partially degraded primer-dimer molecules can serve as templates for a subsequent PCR amplification.

Production of primer-dimers could lower the amplification yield of the desired target region. If primers cannot be selected with dA nucleotides near the ends, consider using primers with 3´ terminal dU-nucleotides. Single-stranded DNA with terminal dU nucleotides are not substrates for UDG (Delort et al., 1985) and, therefore, the primers are not degraded. Biotin-dUMP derivatives are not substrates for UDG.

For more information about designing primers, see "Guidelines for Designing Primers" on page 26.

Do not use UDG in subsequent amplifications of dU-containing PCR template, such as in nested PCR protocols. The UNG degrades the dU-containing PCR product, preventing further amplification.

#### PCR Good Laboratory Practices

When preparing samples for PCR amplification:

- Wear a clean lab coat (not previously worn while handling amplified PCR products or used during sample preparation) and clean gloves.
- Change gloves whenever you suspect that they are contaminated.
- Maintain separate areas and dedicated equipment and supplies for:

Sample preparation

PCR setup

PCR amplification

Analysis of PCR products

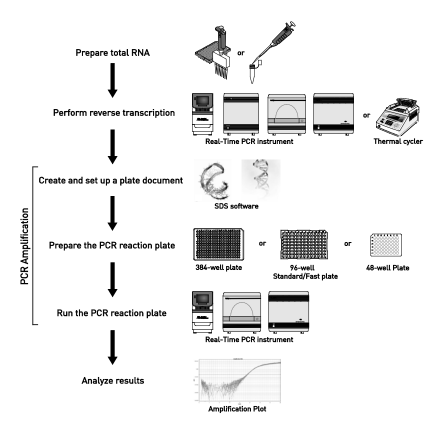
#### PCR Good Laboratory Practices, Continued

- Never bring amplified PCR products into the PCR setup area.
- Open and close all sample tubes carefully. Try not to splash or spray PCR samples.
- Keep reactions and components capped as much as possible.
- Use a positive-displacement pipette or aerosol-resistant pipette tips.
- Clean lab benches and equipment periodically with a 10% bleach solution.

## Methods

### **Procedural Overview**

This diagram is an overview of the procedures for performing gene expression experiments.



### Prepare the Template

#### Examine RNA Template Quality

After isolating the template, examine its quality and quantity and store it properly.

Before using the PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix, you need to synthesize single-stranded cDNA from total RNA or mRNA samples. For optimal performance, the RNA should be:

- Between 0.002 and 0.2 μg/μL
- Less than 0.005% of genomic DNA by weight
- Free of inhibitors of reverse transcription and PCR
- Dissolved in PCR-compatible buffer
- Free of RNase activity

IMPORTANT! If you suspect that the RNA contains RNase activity, add RNase inhibitor to the reverse transcription reaction at a final concentration of 1.0  $U/\mu L$ .

- Nondenatured
- **IMPORTANT!** It is not necessary to denature the RNA. Denaturation of the RNA may reduce the yield of cDNA for some gene targets.

## Examine DNA Template Quality

Use both of the following methods to examine DNA quality:

- Agarose gel electrophoresis Purified DNA should run as a single band on an agarose gel. Agarose gels reveal contaminating DNAs and RNAs, but not proteins.
- Spectrophotometry The A<sub>260</sub>/A<sub>280</sub> ratio should be 1.8 to 2.0. Smaller ratios usually indicate contamination by protein or organic chemicals.
   Spectrophotometry can reveal protein contamination, but not DNA or RNA contamination.

#### Quantitate the Template

Template quantitation is critical for successful PCR reactions. The most common way to determine DNA quantity is to measure the absorbance (optical density or O.D.) of a sample at 260 nm in a spectrophotometer.

One O.D. unit is the amount of a substance dissolved in 1.0 mL that gives an absorbance reading of 1.00 in a spectrophotometer with a 1-cm path length. The wavelength is assumed to be 260 nm unless stated otherwise. A260 values can be converted into  $\mu g/\mu L$  using Beer's Law:

Absorbance (260 nm) = sum of extinction coefficient contributions  $\times$  cuvette pathlength  $\times$  concentration

The following formulas are derived from Beer's Law (Ausubel et al., 1998):

- Concentration of single-stranded DNA =  $A_{260} \times 33 \,\mu\text{g/}\mu\text{L}$
- Concentration of double-stranded DNA =  $A_{260} \times 50 \,\mu\text{g}/\mu\text{L}$
- Concentration of single-stranded RNA =  $A_{260} \times 40 \,\mu\text{g}/\mu\text{L}$

**Note**: Absorbance measurements of highly concentrated (O.D. > 1.0) or very dilute (O.D. < 0.05) DNA or RNA samples can be inaccurate. Dilute or concentrate the DNA/RNA to obtain a reading within the acceptable range.

#### Store the Template

Store the templates as follows:

- Store purified RNA templates at -20°C or -70°C in RNase-free water.
- Store purified DNA templates at -20°C or -70°C in TE, pH 8.0.

### Set Up the Plate Document

## Select a Plate for PCR

Select a plate appropriate for your real-time instrument Refer to page 9 for part numbers of the plates.

#### Configure the Plate Document

For information about configuring plate documents when performing real-time quantification, refer to the appropriate user guides listed in "Support Documents" on page 33.

## Prepare the PCR Reaction Plate

#### General Guidelines

- For best results, it is recommended to perform four replicates of each reaction.
- Reaction mixes can be prepared depending upon your experimental requirements. Scale the components to be included in the reaction mix according to the number of reactions to be performed. Include an additional 10% of the reaction mix volume to account for variations in pipetting.
- If using smaller reaction volumes, scale all components of the reaction mix proportionally. Reaction volumes  $<10~\mu L$  are not recommended.

#### Reminder About Your Primers

Refer to page 26 for information about identifying target sequences and designing primers.

Note: Separate PCR thermal-cycling conditions are required for primers with a  $T_{\rm m}\!<\!60^{\circ}\!C$ 

## Reagent Handling and Preparation

Follow these guidelines to ensure optimal PCR performance. Prior to use:

- Mix the PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix thoroughly by swirling the bottle.
- Place frozen cDNA samples and primers on ice to thaw. After the samples are thawed, vortex them, then centrifuge the tubes briefly.
- **IMPORTANT**: Do not use plastics made of polyethylene terephthalate copolyester, glycol modified (PTEG) for storage of PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix or reaction mixes. SYBR<sup>™</sup> dye has been shown to stick to this type of plastic material.

Plastics recommended for storage include polypropylene, high density polyethylene (HDPE), and polystyrene.

**CAUTION** CHEMICAL HAZARD. PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix (2X) may cause eye, skin and respiratory tract irritation. Read the SDS, and follow the handling instructions. Wear appropriate protective eyewear, clothing, and gloves.

## Prepare the PCR Reactions

1. Prepare the appropriate number of reactions according to the volumes in the following table:

Component	10 μL/well	20 μL/well
PowerUp™ SYBR™ Green Master Mix (2X)	5 μL	10 μL
Forward and Reverse Primers <sup>1</sup>	Variable	Variable
cDNA template + RNase-free water²	Variable	Variable
Total Volume	10 μL	20 μL

 $<sup>^{\</sup>rm I}$  For optimal performance in Fast and Standard modes, use a range from 300–800 nM of each primer.

- 2. Mix the components thoroughly, and centrifuge briefly to spin down the contents and eliminate any air bubbles.
- 3. Transfer the appropriate volume of each reaction to each well of an optical plate.
- 4. Seal the plate with an optical adhesive cover, and centrifuge the plate briefly to spin down the contents and eliminate any air bubbles.

**Note:** PCR can be performed on the reaction plate at any time up to 24 hours after completing the reaction setup when kept at room temperature.

<sup>&</sup>lt;sup>2</sup> For optimal performance, use up to 100 ng of cDNA for each reaction.

### Run the PCR Reaction Plate

Run the plate on an Applied Biosystems real-time quantitative PCR instrument. See the appropriate instrument user guide for help with programming the thermal-cycling conditions or with running the plate.

- 1. Place the reaction plate in the instrument.
- 2. Set the thermal cycling conditions using the default PCR thermal-cycling conditions specified in the following tables according to the instrument cycling parameters and the melting temperature of your primers:

Fast Cycling Mode (Primer T <sub>m</sub> ≥60°C)			
Step	Temperature	Duration	Cycles
UDG Activation	50°C	2 min	Hold
Dual-Lock™ DNA Polymerase	95°C	2 min	Hold
Denature	95°C	1* or 3** sec	40
Anneal/Extend	60°C	30 sec	40

<sup>\*</sup> Denature for 1 second when using ViiA<sup>™</sup>7, or QuantStudio<sup>™</sup> family of Real-Time PCR Systems.

<sup>\*\*</sup>Denature for 3 seconds when using 7500 Fast, StepOne $^{\text{\tiny TM}}$ , or StepOnePlus $^{\text{\tiny TM}}$  Real-Time PCR Systems.

Standard Cycling Mode (Primer T <sub>m</sub> ≥60°C)*			
Step	Temperature	Duration	Cycles
UDG Activation	50°C	2 min	Hold
Dual-Lock <sup>™</sup> DNA Polymerase	95°C	2 min	Hold
Denature	95°C	15 sec	40
Anneal/Extend	60°C	1 min	40

Standard Cycling Mode Primer T <sub>m</sub> <60°C*			
Step	Temperature	Duration	Cycles
UDG Activation	50°C	2 min	Hold
Dual-Lock™ DNA Polymerase	95°C	2 min	Hold
Denature	95°C	15 sec	
Anneal	55-60°C**	15 sec	40
Extend	72°C	1 min	

<sup>\*</sup> For best results, only use the standard cycling mode with 7900HT Real-Time PCR Instruments.

- 3. Set the instrument to perform a default dissociation step.
  - **Note**: A dissociation curve can be performed up to 72 hours after the real-time PCR run if the plate is stored in the dark, or up to 24 hours later if the plate is stored exposed to light.
- 4. Set the reaction volume appropriate for the type of plate being used for your PCR reaction.
- 5. Start the run.

<sup>\*\*</sup>Anneal temperature should be set to the melting point for your primers.

## **Analyze Your Results**

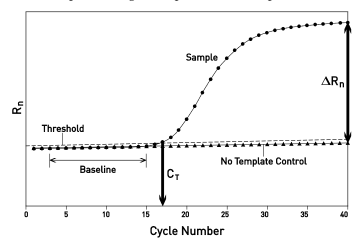
The general process for analyzing the data from gene expression assays requires that you:

- View the amplification plots.
- Adjust the baseline and threshold values to determine the threshold cycles (C<sub>T</sub>) for the amplification curves.
- Use the standard curve method or the relative quantification ( $\Delta\Delta C_T$ ) method to analyze the results.

#### Baseline and Threshold Values

Use the software provided with your instrument to automatically calculate or manually set the baseline and threshold for the amplification curves.

- Baseline refers to the initial cycles of PCR in which there is little change in fluorescence signal.
- The intersection of the threshold with the amplification plot defines the C<sub>T</sub> in real-time PCR assays. The threshold is set above the background and within the exponential growth phase of the amplification curve.

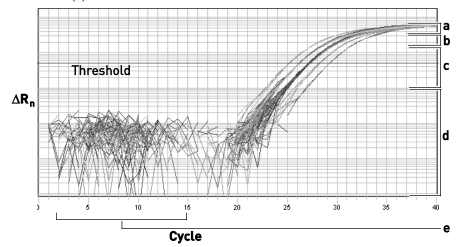


#### View the Amplification Plots

The instrument software calculates baseline and threshold values for a detector based on the assumption that the data exhibit the "typical" amplification curve.

A typical amplification curve, as shown below, has a:

- Plateau phase (a)
- Linear phase (b)
- Exponential (geometric) phase (c)
- Background (d)
- Baseline (e)



#### Manually Adjust the Baseline and Threshold

Experimental error (such as contamination or inaccurate pipetting) can produce data that deviate significantly from data for typical amplification curves. Such atypical data cause the software algorithm to generate incorrect baseline and threshold values for the associated detector.

Reviewing all baseline and threshold values after analysis of the study data is recommended. If necessary, adjust the values manually as described in the appropriate instrument user manual.

**IMPORTANT!** After analysis, you must verify that the baseline and threshold were called correctly for each well by viewing the resulting amplification plots.

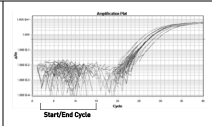
#### **Baseline Settings**

See the example amplification plots below to determine whether or not the baseline and threshold settings were correctly set.

#### **Baseline Set Correctly**

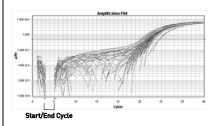
The amplification curve begins after the maximum baseline.

No adjustment necessary.



#### Baseline Set Too Low

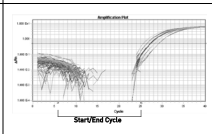
The amplification curve begins too far to the right of the maximum baseline. Increase the End Cycle value.



#### Baseline Set Too High

The amplification curve begins before the maximum baseline.

Decrease the End Cycle value.

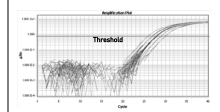


#### Threshold Settings

#### Threshold Set Correctly

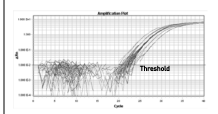
The threshold is set in the exponential phase of the amplification curve.

Threshold settings above or below the optimum increase the standard deviation of the replicate groups.



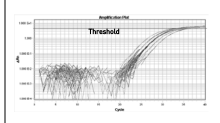
#### Threshold Set Too Low

The threshold is set below the exponential phase of the amplification curve. The standard deviation is significantly higher than that for a plot where the threshold is set correctly. Set the threshold up into the exponential phase of the curve.



#### Threshold Set Too High

The threshold is set above the exponential phase of the amplification curve. The standard deviation is significantly higher than that for a plot where the threshold is set correctly. Set the threshold down into the exponential phase of the curve.



## Analyze the Results

Using the PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix, you can perform two types of quantitation: relative and absolute.

- Relative quantitation compares a target against an internal standard. You can perform relative quantitation using either the standard curve method or the comparative  $C_T$  method.
- Absolute quantitation compares the  $C_T$  of an unknown sample against a standard curve with known copy numbers.

#### Relative Quantitation Method

Gene expression can be measured by the quantitation of cDNA relative to a calibrator sample. The calibrator sample serves as a physiological reference. In a typical experiment, gene expression levels are studied as a function of a treatment of cells in culture, of patients, or of tissue type. The calibrator sample in each case is the cDNA from the untreated cells or patients, or a specific tissue type.

All quantitations are also normalized to an endogenous control (such as GAPDH) to account for variability in the initial concentration and quality of the total RNA, and in the conversion efficiency of the reverse transcription reaction.

### Resources for Data Analysis

For more information about analyzing your data, refer to the appropriate instrument manual available at: **thermofisher.com/lifescience**, or contact Technical Support (see page 33 for a list of support documents).

## **Detect Nonspecific Amplification**

Because SYBR™ Green dye detects any double-stranded DNA, check for nonspecific product formation by using dissociation-curve or gel analysis.

## Dissociation Curves

A dissociation curve is a graph that displays dissociation data from the amplicons of quantitative PCR runs. Change in fluorescence, due to a dye or probe interacting with double-stranded DNA, is plotted against temperature.

When to Generate Dissociation Curves

**Note:** Because of the presence of heat-labile UDG, you can generate a dissociation curve up to 72 hours after the real-time PCR run on any Applied Biosystems Real-Time PCR System.

#### An Example

The dissociation curves below show typical primer-dimer formation. The specific product is shown with a melting temperature ( $T_m$ ) of 80.5°C, but the primer-dimer has a characteristically lower  $T_m$  of 75°C.

Primer-dimers are most prevalent in NTC wells and sample wells containing a low concentration of template.

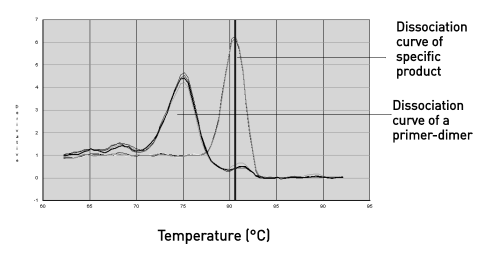


Figure 3 Example of two dissociation curves

(*Optional*) Check PCR Product Purity by Agarose Gel Electophoresis **Note:** Because of the presence of heat-labile UDG, you can verify the absence of nonspecific amplification using agarose gel electrophoresis up to 72 hours after amplification.

1. Load 12 to 15 μL of sample per well on an ethidium bromide-stained agarose gel made with UltraPure<sup>™</sup> Agarose 1000 (Cat. no. 16550-100):

PCR Fragment Size	% Agarose in TBE Buffer	% Agarose in TAE Buffer
<100 bp	5%	6%
100–250 bp	3%	4%

CHEMICAL HAZARD. Ethidium bromide causes eye, skin, and respiratory tract irritation and is a known mutagen (that is, it can change genetic material in a living cell and has the potential to cause cancer). Always use adequate ventilation such as that provided by a fume hood. Read the SDS, and follow the handling instructions. Wear appropriate protective eyewear, clothing, and gloves.

2. Run the gel:

For PCR fragments <100 bp, use 80 to 100 V for 45 to 60 min.

For PCR fragments 100 to 250 bp, use 100 to 115 V for 1 to 1.5 h.

3. Run samples 1/3 to 1/2 the length of the gel, without letting the dye run off the bottom of the gel. Use a UV lamp to check the migration of the samples.

## **Troubleshoot**

Observation	Possible Cause	Action
High C <sub>T</sub> values/poor	Insufficient cDNA template is	Use up to 100 ng of cDNA
precision or failed PCR	present	template per 20-µL reaction.
reactions	Quality of cDNA template is poor	<ul> <li>Quantify the amount of cDNA template.</li> <li>Test the cDNA template for the presence of PCR inhibitors.</li> </ul>
	Sample degradation	Prepare fresh cDNA, then repeat the experiment.
	Incorrect pipetting.	Prepare the reactions as described on page 16.
	Reduced number of PCR cycles in the thermal cycler protocol	Increase the number of PCR cycles to the default setting of 40 (see page 17).
	Primer-dimer formation and residual polymerase activity	<ul> <li>Prepare the reaction mixes and the reaction plate on ice.</li> <li>To ensure optimal results, run the reaction plate as soon as possible after completing the reaction setup. If you cannot run a reaction plate within 2 hours after completing the reaction setup, store the reaction plate at 4°C.</li> </ul>
Low $\Delta R_n$ or $R_n$ values	Extension time is too short	Use the default thermal profile settings (see page 17).
	Primer-dimer formation and residual polymerase activity	<ul> <li>Prepare the reaction mixes and the reaction plate on ice.</li> <li>To ensure optimal results, run the reaction plate as soon as possible after completing the reaction setup. If you cannot run a reaction plate within 2 hours after completing the reaction setup, store the reaction plate at 4°C.</li> </ul>

Observation	Possible Cause	Action	
		1.12.0.2.0	
R <sub>n</sub> vs. Cycle plot is not	ROX <sup>™</sup> dye was not selected as the	Select ROX dye as the passive	
displayed	passive reference when the plate	reference when you set up the	
	document was set up	plate document.	
Extremely high $\Delta R_n$ or $R_n$	ROX dye was not selected as the	Select ROX dye as the passive	
values	passive reference when the plate	reference when you set up the	
	document was set up	plate document.	
	Evaporation	Make sure that the reaction plate	
	•	is sealed completely, especially	
		around the edges.	
Lower $\Delta R_n$ values obtained in	C <sub>T</sub> value is less than 15	Adjust the upper baseline range	
early cycles		to a value less than 15.	
High variability across the	ROX dye was not selected as the	Select ROX dye as the passive	
reaction plate	passive reference when the plate	reference when you set up the	
1	document was set up	plate document.	
	Evaporation	Make sure that the reaction plate	
		is sealed completely, especially	
		around the edges.	
High variability across	Reaction mix was not mixed well	Mix the reaction mix gently by	
replicates		inversion, then centrifuge briefly	
1		before aliquoting to the reaction	
		plate.	
Fluorescent intensity too high	Primer concentration is too high	Use ≤200 nM of each primer.	
(StepOne <sup>™</sup> and StepOnePlus <sup>™</sup>		1	
systems)			
5,5000110)			

## Appendix A

### **Identify Target Sequences and Design Primers**

#### Identify Target Sequence and Amplicon Size

A target template is a DNA sequence, including cDNA, genomic DNA, or plasmid nucleotide sequence that you want to amplify.

Using Primer Express Software, you design primers to amplify amplicons (segments of DNA) within the target sequence. Shorter amplicons work best. Consistent results are obtained for amplicon size ranges from 50 to 150 bp.

#### Guidelines for Designing Primers

Using Primer Express<sup>™</sup> Software

Design primers using Primer Express Software as described in the Primer Express  $^{\text{TM}}$  Version 3.0 Getting Started Guide (PN 4362460) and Online Help.

#### General Guidelines

- Do not overlap primer and probe sequences. The optimal primer length is 20 bases.
- Keep the GC content in the 30–70% range.
- Avoid runs of identical nucleotides. If repeats are present, there must be fewer than four consecutive G residues.
- Make sure the last five nucleotides at the 3' end contain no more than two G and/or C bases.

If the template is	Then
DNA	
plasmid DNA	Design the primers as described above.
genomic DNA	
cDNA	Design the primers as described above. Also see "Select an Amplicon Site for cDNA" on page 27.
RNA	Design the primers as described above.

#### Select an Amplicon Site for cDNA

Selecting a good amplicon site ensures amplification of the target cDNA without co-amplifying the genomic sequence, pseudogenes, and related genes.

#### Guidelines

- The amplicon should span one or more introns to avoid amplification of the target gene in genomic DNA.
- The primer pair must be specific to the target gene; the primer pair does not amplify pseudogenes or other related genes.
- Design primers according to Primer Express Software guidelines.
- Test the amplicons, then select those that have the highest signal-to-noise ratio (that is, low C<sub>T</sub> with cDNA and no amplification with no template control or genomic DNA).
- If no good sequence is found, you may need to examine the sequence and redesign the amplicon or to screen for more sites.

If the gene you are studying does not have introns, then you cannot design an amplicon that amplifies the mRNA sequence without amplifying the genomic sequence. In this case, you may need to run RT minus controls.

## Optimize Primer Concentrations for PCR

#### **Overview**

By independently varying the forward and reverse primer concentrations, you can identify the primer concentrations that provide optimal assay performance. The primer concentrations you select should provide a low  $C_T$  and a high  $\Delta R_n$  when run against the target template, but should not produce nonspecific product formation with NTCs.

#### Quantitate the Primers

- 1. Measure the absorbance (at 260 nm of a 1:100 dilution) of each primer oligonucleotide in TE buffer.
- 2. Calculate the sum of extinction coefficient contributions for each primer:
  - extinction coefficient contribution =  $\Sigma$ (extinction coefficient × number of bases in oligonucleotide sequence)
  - See "An Example Calculation of Primer Concentration" on page 28 for an example calculation.
- 3. Calculate the oligonucleotide concentration in μM for each primer: absorbance at 260 nm = sum of extinction coefficient contribution × cuvette pathlength × concentration/100

  Rearrange to solve for concentration:
  - concentration = 100[absorbance at 260 nm / (sum of extinction coefficient contribution × cuvette pathlength)]

An Example Calculation of Primer Concentration

In this example, the concentration of a primer (in TE buffer, diluted 1:100), with the sequence CGTACTCGTTCGTGCTGC is calculated using the following values:

Chromophore	Extinction Coefficient	Number of Specific	Extinction Coefficient
	Coefficient	Chromophores in Example Sequence	Contribution
Α	15,200	1	15,200
С	7050	6	42,300
G	12,010	5	60,050
T	8400	6	50,400
Total	_	<del></del>	167,950

measured absorbance at 260 nm = 0.13

sum of extinction coefficient =  $167,950 \text{ M}^{-1}\text{cm}^{-1}$  contributions for probe cuvette pathlength = 0.3 cm

Absorbance (260 nm) = sum of extinction coefficient contributions  $\times$  cuvette pathlength  $\times$  oligonucleotide concentration/100

 $0.13 = 167,950 \text{ M}^{-1}\text{cm}^{-1} \times 0.3 \text{ cm} \times \text{C}/100$ 

 $C = 258 \, \mu M$ 

# Determine the Optimal Primer Concentration

WARNING CHEMICAL HAZARD. PowerUp™ SYBR™ Green Master Mix is a combustible liquid and vapor (keep away from heat and flame). It may cause eye, skin, and respiratory tract irritation. Read the SDS, and follow the handling instructions. Wear appropriate protective eyewear, clothing and gloves.

To optimize primer concentrations for PCR:

1. Prepare a 96-well reaction plate as described below.

Use 10 to 100 ng of genomic DNA or 1 to 10 ng of cDNA template.

The final concentration of PowerUp<sup>™</sup> SYBR<sup>™</sup> Green Master Mix is 1X.

**Note**: The plate configuration accounts for four replicates of each of the following nine variations of primer concentration applied to both template and NTC wells:

Reverse Primer	Forward Primer (nM)		
(nM)	300	500	800
300	300/300	500/300	800/300
500	300/500	500/500	800/500
800	300/800	500/800	800/800

2. Calibrate your instrument for SYBR™ Green Dye, if necessary. Refer to the instrument user manual for calibration instructions.

**Note**: It is recommended to calibrate your instrument every 6 months.

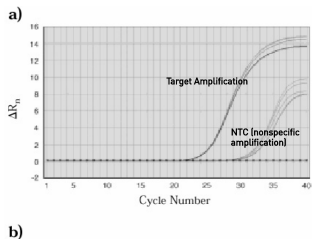
- 3. Load the plate into a Applied Biosystems real-time PCR system.
- 4. Program the thermal-cycling conditions according to the information in step 2 on page 17.
- 5. Run the plate.
- 6. Compile the results for  $\Delta R_n$  and  $C_T$ , then select the minimum forward and reverse primer concentrations that yield the maximum  $\Delta R_n$  values and low  $C_T$  values.

Confirm the Absence of Nonspecific Amplification Dissociation curves help you select the optimal primer concentrations for your  $SYBR^{TM}$  quantification assays.

- Review the linear view of the amplification plot in your NTC wells.
   Note: In Figure A-1 on page 30, part a, the strong amplification of the NTC wells indicates that significant nonspecific amplification is occurring.
- 2. Generate a dissociation curve with your Real-Time PCR System.

**Note**: In the example dissociation curve data shown in Figure A-1 on page 30, part b, the melting temperature of the product generated in the absence of template is lower than the melting temperature of the specific product generated with template. This variation is typical of primer-dimer formation, and it indicates that lower primer concentration may provide optimal results.

#### Example of Nonspecific Amplification



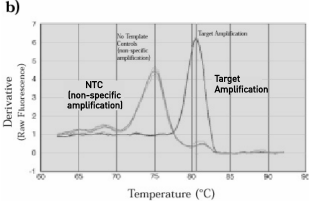


Figure A-1 Amplification data using SYBR™ Green dye chemistry

- (a) Amplification plot (linear view) demonstrating suspected nonspecific amplification in NTC wells.
- (b) Dissociation curve analysis confirming that product in NTC wells has a melting temperature different from the specific product.

## Appendix B

## Safety

## Chemical Safety Guidelines

To minimize the hazards of chemicals:

- Read and understand the Safety Data Sheets (SDSs) provided by the chemical manufacturer before you store, handle, or work with any chemicals or hazardous materials.
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing). For additional safety guidelines, consult the SDS.
- Minimize the inhalation of chemicals. Do not leave chemical containers open. Use only with adequate ventilation (for example, fume hood). For additional safety guidelines, consult the SDS.
- Check regularly for chemical leaks or spills. If a leak or spill occurs, follow the manufacturer's cleanup procedures as recommended in the SDS.
- Comply with all local, state/provincial, or national laws and regulations related to chemical storage, handling, and disposal.

#### Chemical Waste Safety Guidelines

To minimize the hazards of chemical waste:

- Read and understand the Safety Data Sheets (SDSs) provided by the manufacturers of the chemicals in the waste container before you store, handle, or dispose of chemical waste.
- Provide primary and secondary waste containers. (A primary waste container holds the immediate waste. A secondary container contains spills or leaks from the primary container. Both containers must be compatible with the waste material and meet federal, state, and local requirements for container storage.)
- Minimize contact with chemicals. Wear appropriate personal protective equipment when handling chemicals (for example, safety glasses, gloves, or protective clothing). For additional safety guidelines, consult the SDS.
- Minimize the inhalation of chemicals. Do not leave chemical containers open. Use only with adequate ventilation (for example, fume hood). For additional safety guidelines, consult the SDS.
- Handle chemical wastes in a fume hood.
- After emptying the waste container, seal it with the cap provided.
- Dispose of the contents of the waste tray and waste bottle in accordance with good laboratory practices and local, state/provincial, or national environmental and health regulations.

#### Waste Disposal

If potentially hazardous waste is generated when you operate the instrument, you must:

- Characterize (by analysis if necessary) the waste generated by the particular applications, reagents, and substrates used in your laboratory.
- Ensure the health and safety of all personnel in your laboratory.
- Ensure that the instrument waste is stored, transferred, transported, and disposed of according to all local, state/provincial, and/or national regulations.

**IMPORTANT!** Radioactive or biohazardous materials may require special handling, and disposal limitations may apply.

#### Biological Hazard Safety

WARNING BIOHAZARD. Biological samples such as tissues, body fluids, infectious agents, and blood of humans and other animals have the potential to transmit infectious diseases. Follow all applicable local, state/provincial, and/or national regulations. Wear appropriate protective equipment, which includes but is not limited to: protective eyewear, face shield, clothing/lab coat, and gloves. All work should be conducted in properly equipped facilities using the appropriate safety equipment (for example, physical containment devices). Individuals should be trained according to applicable regulatory and company/institution requirements before working with potentially infectious materials. Read and follow the applicable guidelines and/or regulatory requirements in the following:

- U.S. Department of Health and Human Services guidelines published in Biosafety in Microbiological and Biomedical Laboratories
   http://bmbl.od.nih.gov
- Occupational Safety and Health Standards, Bloodborne Pathogens (29 CFR§1910.1030;
  - $http://www.access.gpo.gov/nara/cfr/waisidx\_01/29cfr1910a\_01.html).\\$
- Your company's/institution's Biosafety Program protocols for working with/handling potentially infectious materials.

Additional information about biohazard guidelines is available at: http://www.cdc.gov

## **Documentation and Support**

#### Support Documents

You can download the following documents from: thermofisher.com/lifescience

Document	Part number
All Systems	
High-Capacity cDNA Reverse Transcription Kit Protocol	4375575
Primer Express™ Software Version 3.0 Getting Started Guide	4362460
Real-Time PCR Systems Chemistry Guide	4348358
StepOne and StepOnePlus Systems	
Applied Biosystems StepOne™ Real-Time PCR System Getting Started Guide for Relative Standard Curve and Comparative Cτ (ΔΔCτ) Experiments	4376785
Applied Biosystems StepOne™ Real-Time PCR System Getting Started Guide for Standard Curve Experiments	4376784
Applied Biosystems StepOne™ Real-Time PCR System Installation, Maintenance, and Networking Guide	4376782
7500 Fast System	
Applied Biosystems 7300/7500/7500 Fast Real-Time PCR System Installation and Maintenance Getting Started Guide	4347828
Applied Biosystems 7300/7500/7500 Fast Real-Time PCR System Relative Quantification Getting Started Guide	4347824
Applied Biosystems 7300/7500/7500 Fast Real-Time PCR System Absolute Quantification Getting Started Guide	4347825
7900HT System	
Applied Biosystems 7900HT Fast Real-Time PCR System and SDS Enterprise Database User Guide	4351684
Applied Biosystems 7900HT Fast Real-Time PCR System User Bulletin: Performing Fast Gene Quantification	4352533
ViiA <sup>™</sup> 7 System	
Applied Biosystems ViiA™ 7 Real-Time PCR System Getting Started Guides	4441434
Applied Biosystems ViiA™ 7 Real-Time PCR System User Guide	4442661
QuantStudio™ 12K System	
Applied Biosystems QuantStudio™ 12K Real-Time PCR System User Guide	4470050

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- Submit a question directly to Technical Support (techsupport@lifetech.com)
- Search for user documents, SDSs, vector maps and sequences, application notes, formulations, handbooks, certificates of analysis, citations, and other product support documents
- Obtain information about customer training
- Download software updates and patches

(SDS)

Certificate of **Analysis** 

The Certificate of Analysis provides detailed quality control and product qualification information for each product. Certificates of Analysis are available at thermofisher.com/techresources/support

Limited warranty

Life Technologies and/or its affiliate(s) warrant their products as set forth in the Life Technologies General Terms and Conditions of Sale found on the Life Technologies website at http://www.lifetechnologies.com/termsandconditions. If you have any questions, please contact Life Technologies.

